PATENT SPECIFICATION

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We, TEROSON GMBH, a German Body Corporate, of Hans-bunte-(71)Strasse 4, 6900 Heidelberg 1, Germany, do hereby declare the invention for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:-

The invention relates to novel plastisols containing acrylate polymers and organic

plasticisers.

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Plastisols are generally understood to mean dispersions of organic plastics in plasticisers, which are gelled on heating to an elevated temperature. Conventional plastisols generally comprise pulverulent polyvinyl chloride dispersed in a compatible liquid plasticiser which forms a paste. Polyvinyl chloride plastisols are used for numerous different purposes, particularly as sealing compounds, as anti-corrosive coatings for metals, for impregnating and coating textile substrates and as cable insulations.

However, several serious problems occur in the production and use of PVCplastisols. In fact, the production of PVC itself represents a problem because persons working in the production area may be exposed to a health risk by the vinyl chloride. Of late this has led to the closing of a large number of PVC factories. Residual monomer in the PVC could also endanger people during processing, and possibly even the final consumer if the plastigel comes into contact with foodstuffs.

PVC-plastisols also suffer from the disadvantages in that PVC is both sensitive to light and heat, and that there is a tendency to split of hydrogen chloride. The splitting off of hydrogen chloride represents a particularly serious problem because during use the plastisol must be heated to an elevated temperature. Any hydrogen chloride liberated under these conditions has a corrosive effect and, for example, corrodes metal substrates. Attempts have been made to counteract this thermal decomposition by adding heat stabilisers, but these compounds are frequently toxic so that they cannot be used where the plastisol comes into contact with foodstuffs. These problems are further exacerbated when, in certain species cases, PVC-plastisols require relatively high baking temperatures in order to obtain very short gelling times. In

these circumstances the danger of PVC-decomposition is further increased. According to the present invention, there is provided a plastisol, comprising (a) 100 weight parts of solid particles of average size 0.1 to 500 microns of an organic

polymer consisting substantially of units derived from

(i) one or more of: t-butyl acrylate, a C1-4 alkyl methacrylate or cyclohexyl

methacrylate, or (ii) one or more of the above monomers and one or more comonomers selected from methacrylates of aliphatic C2 to C10 alcohols, acrylates of aliphatic C1-C10 alcohols, styrene and alpha-methyl styrene, said polymer having a glass transition



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temperature of above 35°C and an average degree of polymerisation of more than 400, and (b) 30 to 1000 weight parts of an organic liquid component which serves as a compatible plasticiser for the organic polymer, to form a viscous dispersion having a liquid or paste-like appearance and a viscosity index V_{8/0} (as hereinafter defined) of less than 3.0.

These plasticals possess the excellent processing and product qualities of PVC-

These plastisols possess the excellent processing and product qualities of PVC-plastisols, but gel at lower temperatures and are also chlorine-free so that there can be no splitting off of the chlorine or hydrogen chloride. The average particle size of the polymer used is preferably increased proportionally with the decrease of its glass transition temperature T_g. The glass transition temperature is preferably above 60°C.

To obtain usable results, the degree of polymerisation of the polymer used must be at least 400. Advantageously the degree of polymerisation is between 400 and 20,000, corresponding to an average molecular weight of about 40,000 to 2,000,000.

The preferred polymer component consists substantially wholly of one kind of units derived from t-butyl acrylate, methyl, ethyl, n-propyl, isopropyl, n-butyl, sec-butyl or t-butyl methacrylate or cyclohexyl methacrylate, i.e. is essentially a homopolymer. Methyl methacrylate is preferred. The alcohol acrylate or methacrylate comonomers preferably have an alkanol alcohol component, i.e. are unsubstituted and saturated, and the type (i) monomers, i.e. those listed above, will generally predominate over those of type (ii). The preferred copolymers are those in which one of the monomers is methyl methacrylate and constitutes at least 40%, preferably at least 50%, by weight of the total monomer units, and the other component is a C_{2-4} alkyl methacrylate, especially n-butyl methacrylate, or a C_{1-4} alkyl acrylate.

However, only those copolymers whose glass transition or solidification temperature is above 35°C can be employed in the plastisols of the present invention. The glass transition temperature T_g of a particular copolymer can be calculated beforehand by the Fox formula (T. G. Fox, Bull. Am. Phys. Soc., vol. 1, 123 (1956)):

$$\frac{1}{T_{c_0}} \frac{W_1}{T_1} + \frac{W_2}{T_2} + \cdots + \frac{W_n}{T_n}$$

wherein W_1 represents the weight portion of monomer 1, W_2 the weight portion of monomer 2, T_1 the glass transition temperature of the polymerised monomer 1 in ${}^{\circ}K$, T_2 the glass transition temperature of the polymerised monomer 2 in ${}^{\circ}K$, T_{cn} the glass transition temperature of the copolymer in ${}^{\circ}K$.

The average particle size of the polymer used must be from 0.1 µm to 500 µm, preferably from 0.3 to 200 µm. If too large a particle size is used, the plastisol obtained will not have an adequate internal bond, and no complete gelling of the polymer occurs. If the particle size is too small, the storage stability is inadequate, i.e. gelling occurs even when the polymer is left to stand.

The accompanying drawing is a graph in which the logarithm of the average particle size (in µm) of the organic polymer (a) has been plotted against the glass transition temperature T_g (in °C) of this polymer. There is an interdependence between the average particle size of the polymer (a) and its glass transition temperature T_g such that the polymers most suitable for the plastisols according to the invention substantially occur within the triangle XYZ formed by the coordinates (minimum glass transition temperature, log [maximum average particle size]), (105°C, log [maximum average particle size]) and (105°C, log [minimum average particle size]). It can also be seen that the higher the glass transition temperature of the polymer, the lower can be its average particle size. However, extremely fine-grained polymers with a low glass transition temperature are not suitable.

It is also within the scope of the present invention to use two polymers having different average particle sizes. As a result of using a mixture of a fine-grained and a coarse-grained polymer, it is possible to control with certain limits the flow characteristics and plasticiser requirements (cf. e.g. German Patent Specification No. 934,498).

Both suspension-polymerized and emulsion-polymerized acrylic polymers are suitable for the plastisols according to the invention. In the case of suspension-polymerisation, the desired particle size can be controlled by the stirring rate, and in the case of emulsion-polymerisation by the quantity and type of emulsifier. The particle size of the polymer is determined in conventional manner, e.g. by means of a Coulter counter.

Preferably 65 to 800 parts by weight of plasticiser are used per 100 parts by weight of polymer.

The plasticiser selected must of course be completely compatible with the particular acrylic polymer. A useful simple test of compatibility is that at last 8 days after the gelling of the plasticiser, the plasticiser does not exude, as shown by a completely dry and non-tacky surface. Naturally, the compatibility is dependent both on the type of acrylic polymer and on the type of plasticiser. For example, dibutylphthalate is not a suitable plasticiser for polymethylmethacrylate because compatibility is inadequate. However, the same plasticiser can be used for copolymers of methylmethacrylate and approximately 5 to 25% of butyl methacrylate.

A further important criterion for selecting suitable plasticisers is the storage stability of the plastisol. This can be quantitatively defined by the viscosity index (based

e.g. on a 1:1 polymer/plasticiser mixture) in the following manner:

$$V_{8/0} = \frac{\eta_{8d}}{\eta_0}$$

wherein $V_{8/0}$ represents the viscosity index η_0 the initial viscosity of the plastisol, and

 $\eta_{\rm sd}$ the plastisol viscosity after standing for 8 days.

The viscosity is determined by means of conventional methods of determination, e.g. with a rotary viscosimeter (Drage). According to the definition indicated above the viscosity index of the plastisol of the present invention must be below 3.0, i.e. the viscosity may only rise a maximum of less than three times the initial viscosity of the plastisol within a period of 8 days. Such a plastisol would still not necessarily always be adequately stable, but this can easily be remedied, e.g. by increasing the plasticiser content or adding fillers or other additives.

The selection of the most appropriate plasticiser can again be explained with reference to the accompanying drawing. Within the triangle XYZ, which encloses the suitable acrylic polymers, there is a smaller triangle for each usable plasticiser within which are located those acrylic polymers which can be combined with a particular plasticizer to give a usable plastisol. The zones for the individual plasticisers are limited to the left by the compatibility with the polymer, and upwards (hypotenuse of the triangle) by the viscosity index (storage stability). For each plasticiser these limits can be determined by simple experiments. As examples, the drawings show the zones GYH, DEF and ABC for dimethoxy-ethylphthalate (DMEP), dibutylphthalate (DBP) and dioctylphthalate (DOP).

Plasticisers having at least two aromatic rings and/or two ether groups in the molecule have proved particularly suitable for polymethyl methacrylate. The first group includes, for example, butylbenzyl phthalate, dibenzyl toluene, dibenzyl phthalate, diphenyloctyl phosphate, triphenyl phosphate, tricresyl phosphate, dibenzyl benzoate, and diphenylether. The second group includes, among others, dimethoxy-ethyl phthalate, diethoxyethyl phthalate, dibutoxyethyl phthalate and methylphthalylethyl glycollate. Both criteria are satisfied by diethyleneglycol dibenzoate and dipropyleneglycol di-

benzoate. A further preferred plasticiser is acetyl tributyl citrate.

When methacrylic copolymers are used, generally the compatibility with plasticisers increases, but in many cases there is a viscosity index of at least 3.0, i.e. an inadequate stability. For example, for a copolymer of 15% n-butyl-methacrylate and 85% methyl methacrylate with an average particle size of $100~\mu m$, dibutyl phthalate, diisobutyl phthalate and diamyl phthalate are also suitable in addition to the above-indicated plasticisers. However, if in the case of these polymers the particle size is reduced to $2~\mu m$, the viscosity index for most of the indicated plasticisers rises to at least 3.0. However, tricresyl phosphate and dibenzyl toluene still remain suitable as plasticisers.

- Conventional plasticisers such as dioctyl phthalate, dinonyl phthalate and others require a still higher comonomer content in the copolymer in order to obtain an adequate

compatibility.

In the case of a copolymer consisting of 50 parts by weight of butyl methacrylate and 50 parts by weight of methyl methacrylate and having an average particle size of 50 µm, even if the necessary compatibility exists, it is impossible to obtain an adequate storage stability with most of the above-indicated plasticisers. However, dioctyl phthalate, dinonyl phthalate, didecyl phthalate, butyl cyclohexyl phthalate and the like remain suitable as plasticisers in such cases. Thus, with an increasing comomomer content in the polymer, the aliphatic groups in the phthalate may proportionally become longer. While pure aliphatic plasticisers such as adipates or sebacates are unsuitable it is then possible to use higher boiling aromatic, hydrocarbons with aliphatic groups. For example, Ingralur 839 (hydrocarbon mixture containing more than 40% aromatics, produced by the Fuchs Company, Mannheim) has proved suitable for the

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$$R_1$$
 $CH_2 = C - (CH_2)_n - R_2$

Preferably, a heterocyclic vinyl compound of the general formula

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is used as a comonomer for the co-polymerisation, in which n is 0 or 1, R_1 is a hydrogen atom, a linear or branched alkyl group of from 1 to 4 carbon atoms or a phenyl group and R_2 is a group of the formula

$$O$$
 \parallel
 $-C$
 OR_3 , $-COR_3$, or $-R_3$,

in which R₃ is a heterocyclic group having one or several rings with at least one nitrogen atom in the ring.

Suitable monomers are, for example N- or C-vinyl or N- or C-allyl compounds of imidazole, imidazoline, imidazolidine, benzimidazole, triazole, pyrrole, pyrazole, oxyzole, pyridine, chinoline, diazine as well as other heterocyclic compounds with at least one nitrogen atom in the ring system, N-vinyl imidazole is particularly preferred. Furthermore, vinyl and allyl ethers and acrylic or methacrylic acid esters of the heterocyclic compounds are useful.

The adhesion of the plastisols can be further improved by adding additional cross-linking agents which react with the heterocyclic group of the co-polymerised monomer. Preferably the plastisol contains the cross-linking agent in an amount of from 0.1 to 3.0% by weight. Epoxy resins were found suitable; apart from the commercial epoxy resins a polymer can also be used which, according to the present invention has been co-polymerized with a small amount of acrylate and/or methacrylate monomer with epoxy groups. Furthermore, polybasic carboxylic acids, e.g. benzene tricarboxylic acid, adipic acid, maleic acid and itaconic acid, have proved to be suitable, either as the only cross-linking agent present or together with a polyepoxide, to effect a further improvement of the adhesion properties.

Preparation of the plastisols according to the present invention can also be effected by blending together two or more acrylate or methacrylate polymers which meet the requirements of the invention. For example, a methyl methacrylate homopolymer can be blended with a methyl methacrylate copolymer or with a copolymer which has been further modified by co-polymerisation with a heterocyclic compound. In this manner, particularly favourable rheological properties can be obtained.

It is also possible to blend copolymers containing groups which react with each other. The above-described combination of copolymers with epoxy groups and of copolymers with heterocyclic groups is one example for this method.

The plastisols according to the invention can be processed in conventional manner dependent on the particular viscosity, for example by applying with a float, coating with a brush, applying with a paint gun, spraying with an atomiser and air, spraying by high pressure (without air), scraping on, calendering, pouring or immersing.

Finally, to form the plastigel the plastisols must be gelled. For this purpose temperatures between about 70 and 240°C are used, dependent on the plastisol composition and the particular working conditions. The necessary gelling time is between two minutes and two hours. The necessary heating can take place, for example, by hot air, metal pipes, infra-red rays, high frequency heating, etc.

The plastisols according to the invention can be shaped and gelled to form shaped plastics articles. The plastisols are particularly suitable as abrasion and corrosion inhibitors for steel sheets, for example as underbody-protection for automobiles, trucks and buses. The plastisols can also be used for seam seals for levelling out folding systems, as metal adhesives, coating compounds, spot welding compounds, expanding pastes, sealing agents for spot welded sheets and for sealing double-flanged seams. They can also be used for impregnating and coating textiles and similar substrates such as tarpaulins, artificial leather, floor coverings, carpet backings, packaging materials, conveyor belts and driving belts. Using the immersion process, it is possible to produce hollow bodies, gloves, boots, finger stalls etc. The plastisols may also be shaped and gelled to form sealing gaskets, such as seals for bottle caps and folding seams, as plastic seals for air and oil filters, for extrusion and injection moulding highly elastic sections and items such as sealing strips and packaging cords, trays, technical components and for rolling foils and strips. If appropriate plasticisers and additives are used, the plastisols according to the invention are also suitable for packaging foodstuffs because there are no objections to methyl methacrylate.

The following Examples 3, 4, 6, 8, 10, 15 to 20 and 25 to 34 illustrate the present invention. Examples 1, 2, 5, 7, 9, 11 to 14, and 21 to 24 are comparative. All formulations are given in parts by weight unless otherwise stated.

Examples 1 to 15.

Table 1 gives the results of 15 Examples. In Examples 1, 2, 7, 11 and 12 the

plasticiser used was unsuitable for the particular polymer due to inadequate compatibility. In Examples 5, 9, 13 and 14 the value of 3.0 for the viscosity number was exceeded when using the indicated plasticiser, so that these combinations are also unsuitable. However, excellent results were obtained with the polymer/plasticiser combinations of Examples 3, 4, 6, 8, 10 and 15, wherein the plasticiser satisfied was both completely compatible with the polymer and the viscosity index V_{8/0} of the resulting plastisol was less than 3.0.

	, d	Ę		. :		Viscosity in cP	in cP	14	
Example	Farts by weight of nolvmei	go ii oʻ	Particle diameter in <i>u</i> m	Parts by weight of plasticiser	Compatibility	immediatelv	after 8 days	, °, °, °, °, °, °, °, °, °, °, °, °, °,	Elasticity
1	SO DAMA	105	30	50 DOP	had	910	066	1.09	good
4	TANATA TO CO)) !		i I	
7	6	**		50 DIBP	moderate	720	1180	1.64	good
ec.	6	66	. 66	50 TKP	good	2630	3070	1.17	very good
4	,	66	66	50 DPGB	good	3450	3770	1.01	very good
	50 MMA/ BMA 85/15	68	7	50 DIPB	good	2900	solid	. 8	good
9	•	6	•	50 TKP	poog	9050	21000	2.32	very good
7	6	2	"	śo dop	moderate	4850	7300	1.51	good
∞			· 66	50 DBT	good	2870	6850	2,39	very good
6	50 MMA/ BMA 50/50	9.5	50	50 DIBP	. pood	8200	solid	8	very good
10	•	Ç	66	50 DOP	good	15900	35000	2.20	very good
11	6		. 6	50 DOA	moderate	1150	2400	2.09	good
12		ç	°	50 TEHM	moderate	13600	13400	86.0	hard, elastic
13	50 MMA/ BMA 30/70	40	240	50 DIBP	poog	5250	solid	8	soft, tacky
14				50 DOP	good	32750	solid,	8	very good
15	,			50 TEHM	good	18100	14000	. 22.0	pood
BMA – PMMA – PMMA – PMMA –	n-butylmethacrylate dioctylphthalate polymethyl methacry	nacrylate lalate methacrylate		DPGB – dip DBT – dib DOA – dio TFHM – tri(dipropylene glycoldibenzoate dibenzyltoluene dioctyl adipate tri(ethylbexyl)mellitate	oenzoate ate		-	

diisobutyl phthalatetricresyl phosphate DIBP TKP

tri(ethylhexyl)mellitate Methyl methacrylate TEHM -

		-
	Example 16. The following mixture is homogenised with a stirrer and applied with a doctor blade to a degreased steel sheet:	
	30 parts PMMA (30 μm)	
5	30 parts butylbenzyl phthalate	5.
3	36 parts chalk	,
	3 parts triethyleneglycol dimethacrylate	
	0.3 parts tertbutylperbenzoate	•
	If this mixture is gelled for 20 minutes at 170°C, a tough strongly adhering film	-
10	is obtained which has good anti-corrosive properties.	10
10	is obtained which has good and-corrosive properties.	
	Example 17.	
	30 parts copolymer (MMA/BMA 50/50) (50 µm)	-
		•
	30 parts dinonyl phthalate	
1.5	40 parts chalk	15
15	5 parts resorcinol resin (Rousselot RH 74—01)	15
	This mixture was tested as a metal adhesive by gelling it at 170°C between two	
	metal plates of (a) steel and (b) aluminium. The subsequent tearing revealed a tensile	
	strength of (a) 52 kg/cm ² and (b) 42 kg/cm ² (5 cm ² bonding area and 0.3 mm	
20	coating thickness). However, if in place of the 50 µm copolymer, a copolymer with	•
20	an average particle size of 500 μ m is used, a tearing value of only 3 kg/cm ² is obtained.	20
	Empreson 1 o 10	
	Example 18.	
	24 parts MMA/BMA 85/15 (2 μm)	•
25	33 parts dibenzyl toluene	
25	40 parts chalk	25
	3 parts azodicarbonamide	
	This mixture gelled in 15 minutes at 170°C and formed a highly elastic solid	
	foam with 100% foam expansion.	
	Emana mia 10	-
20	Example 19.	20
30	20 parts PMMA (particle size 30 μm)	30
	30 parts butylbenzyl phthalate	
	45 parts chalk	
	5 parts resorcinol resin (Rousselot RH 74—01)	,
	This highly viscous substance was suitable as a sealing compound for metal welding	25
35	seams in the automobile and refrigerator industry. After baking at 180°C for 15 minutes	:35
	the substance adhered well to polished metal sheets and can be easily lacquered.	
	T1- 20	
	Example 20.	
	If a polymer plasticiser (a liquid urea/formaldehyde plasticiser resin sold as	
4.0	Plastigen) is used the following comparison is obtained with a polymer:plasticiser	40
40	mixing ratio of 1:1.	40

TABLE 2

	Dantista	Viscosity	in cP			
Polymer	Particle size µm	immediately	after 8 days	Compati- bility	Elasticity	
100% PMMA	30	22000	16000	good	hard	
MMA/BMA 85/15	2	163000	218000	good	hard, tough	
MMA/BMA 50/50	60	23250	22250	good	elastic	

Example 21 (comparison).

Table 3 shows the thermal stability of an acrylic plastisol as compared with a PVC-plastisol with a baking cycle of 30 minutes at 150°C.

TABLE 3

	· ·		1	2 .	
	Emulsion PVC, K-value	74 (2 μm)	20		
	PMMA (30 μm)			20	
	Chalk		45	45	
	Butylbenzylphthalate	-	35	35	
	Viscosity in cP	immediately after 10 days	36000 21000	52500 18000	
		V _{10/0}	0.58	0.34	
	Cold elasticity	-40°C	++	++	
	Discolouration after 14 days at	120°C 150°C	slight black	slight slight	
	Abrasion (mm/min) after 14 days storage at	amb. temp. 120°C 150°C	0.016 0.039 1.280	0.045 0.390 0.220	
	the acrylic plastigel the abra rose by a maximum of 5 times		not originally es	specially good, only	
	On comparing the follow:	•		l u m)	
	25 parts PVC (paste typ 40 parts chalk 5 parts iron oxide (mag 27 parts DOP	ing mixtures: `ee, emulsion-polymerisegn.)		ļum)	
-	25 parts PVC (paste type 40 parts chalk 5 parts iron oxide (magaziron parts DOP 3 parts triethyleneglyco 0.3 parts tert-butylperbeat and the same mixture with	ing mixtures: `e, emulsion-polymerise gn.) ol dimethacrylate nzoate	ed, K value 74, 2		
	25 parts PVC (paste typ 40 parts chalk 5 parts iron oxide (mag 27 parts DOP 3 parts triethyleneglyco 0.3 parts tert-butylperber	ing mixtures: ce, emulsion-polymerise gn.) cl dimethacrylate nzoate IA/BMA 50/50) (50 hem between two over ing, in the case of the nilst when left standing ecomposition took pla	um particle diamed apping steel she in air there is property and the compositions are the compositions.	meter) ets and then joining sition and hydrogen ronounced corrosion	
	25 parts PVC (paste type 40 parts chalk 5 parts iron oxide (mag 27 parts DOP 3 parts triethyleneglycon 0.3 parts tert-butylperber and the same mixture with 25 parts copolymer MM in place of PVC by coating the two sheets by spot welding the two sheets by spot welding the two sheets by spot welding the the points where PVC-dicase of the methacrylate plastic On gelling four plastisols a) 60 parts PVC (as in Example 1) 60 parts PVC (as in Example 2) and parts PVC (as in Example 3) and parts PVC (as in Example 4) and parts PVC (as in Example	ing mixtures: be, emulsion-polymerise gn.) ol dimethacrylate nzoate IA/BMA 50/50) (50 hem between two over ing, in the case of the nilst when left standing ecomposition took pla isol. Example 23 (comparise	um particle diameter and the state of the st	meter) ets and then joining sition and hydrogen ronounced corrosion	
	25 parts PVC (paste type 40 parts chalk 5 parts iron oxide (mag 27 parts DOP 3 parts triethyleneglycon 0.3 parts tert-butylperber and the same mixture with 25 parts copolymer MM in place of PVC by coating the two sheets by spot welding the two sheets by spot welding the two sheets by spot welding the points where PVC-dicase of the methacrylate plastic On gelling four plastisols	ing mixtures: be, emulsion-polymerise gn.) ol dimethacrylate nzoate IA/BMA 50/50) (50 hem between two over ing, in the case of the nilst when left standing ecomposition took pla isol. Example 23 (compari iple 22) hple 22)	um particle diameter and the state of the st	meter) ets and then joining sition and hydrogen ronounced corrosion	

			Polymer Particle size in microns	10	10	ν	10
n the	was used unctional stoperties attached 15		Substrate	degreased metal sheet	primed metal sheet	untreated metal sheet	Smoothened Metal 'sheet
On gelling together the following components 30 parts PVC (as in Example 22) 30 parts DOP 20 parts chalk 20 parts barite 30 parts MMA/BMA 85/15 (2 µm) in place of PVC otherwise as a) 30 minutes at 70°C, in the case of a) a crumbly substance is obtained and in the of b) a solid tough film.			Bonding Strength in kg/cm ²	25	4,2	7,2	Film cannot be removed
Example 24 (comparison). Ilowing components ple 22) 15 (2 \(\mu \mu \m) in place of PVC recase of a) a crumbly sub-	Examples 25 to 28. following plastisols in each by weight of a methacryla this manner were tested Example 17. The resultion values were obtained.	TABLE 4	Parts by weight Filler		35 chalk		
On gelling together the following components 30 parts PVC (as in Example 22) 30 parts DOP 20 parts chalk 20 parts barite b) 30 parts barite b) 30 parts MMA/BMA 85/15 (2 µm) in place of or 30 minutes at 70°C, in the case of a) a crur case of b) a solid tough film.	Examples 25 to 28. For the preparation of the following plastisols in each case a copolymer which was polymerised with 1% by weight of a methacrylate monomer with a forup. The plastisols obtained in this manner were tested for their adhesion pusing the method described in Example 17. The results compiled in the Table 4 show that excellent adhesion values were obtained.		Parts by weight Plasticizer	50 DBT	45 DBT	50 DBT	50 DBT
On a) 30 pg 30 pg 30 pg 5 20 pg 5 20 pg 6 pg for 30 pg for 30 pg 6 pg	10 For which will group. To using the 15 Table 4 s		Parts by Weight Polymer	50 MMA/BMA/ methacrylic acid (84:15:1)	20 MMA/BMA/ methacrylic acid (84::15:1)	50 MMA/BMA/ glycidyl methacrylate acrylate (84:15:1)	50 MMA/BMA/ dimethyl amino ethyl methacrylate (84:15:1)
			Example	25	26	27	

10 1,516,510 10 Example 29. (a) A plastisol was prepared from 50 parts by weight of a copolymer from methylmethacrylate and n-butylmethacrylate (MMA/BMA 85:15; particle size approximately 2 μ m) and 50 parts of dibenzyl toluene. 5 This pastisol was tested as metal-to-metal adhesive by applying it to two degreased 5 metal sheets (bonding area 5 cm² and 0.3 mm layer thickness) and gelling it for 30 minutes at 180°C. The subsequent tear-off test (tearing speed: 100 mm per minute) did not yield a bonding strength value. (b) The test was repeated using 50 parts by weight of a terpolymer which was 10 copolymerised with 1 part by weight of N-vinyl imidazole (MMA/BMA/vinyl 10 imidazole 84:15:1). Under otherwise unaltered conditions a bonding strength value of 13 kg/cm² was now obtained. When the above plastisol, with a content of N-vinyl imidazole, according to the invention was used for the bonding of test sheets which had been electro-coated with 15 a dip primer as it is used in the automotive industry a bonding strength value of 15 28 kg/cm² was obtained. Example 30. A further plastisol was prepared from 20 parts by weight of copolymer (MMA/BMA/N-vinyl imidazole 84:15:1) of particle size 10 microns, 35 parts by weight of chalk and 45 parts by weight of dibenzyl toluene. Upon gelling for 30 minutes 20 20 at 180°C on an EC primed metal sheet a bonding strength value of 11 kg/cm² was obtained. The same plastisol surprisingly yielded bonding strength values of 14 to 12 kg/cm², respectively, when applied to degreased or slightly greasy metal sheets. 25 If, for comparative purposes, a methyl methacrylate homopolymer or a methyl 25 methacrylate/butyl methacrylate copolymer is used which was not copolymerised with an adhesion promoter, no bonding strength value was obtained. Example 31. A plastisol was prepared from 50 parts by weight of a copolymer (MMA/N-vinyl imidazole 99:1) of particle size 20 microns, 30 parts by weight of chalk and 45 parts 30 30 by weight of butylbenzyl phthalate. Upon gelling for 30 minutes at 90°C the plastisol coating on the primed metal sheet could not be removed. If, however, the imidazole group is left out of the polymer no adhesion is obtained.

Example 32.

Following Example 29 (b) a plastisol was prepared to which 1% by weight of one of the polybasic carboxylic acids listed below was added. The results show that the adhesion was considerably improved.

TABLE 5 Bonding Strength Values in kg/cm² after Gelling for 30 Minutes at

Acid	120°	160°
None	11.5	20.0
Benzene tricarboxylic acid	20.2	28.0
Adipic acid	22.3	19.6
Maleic acid	34.2	30.7
Itaconic acid	27.8	25.3

Example 33.

A plastisol was prepared from the following ingredients:

100 parts poly(methyl methacrylate), of 35 microns particle size

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⁸⁵ parts acetyl tributyl citrate

⁵ parts titanium dioxide

⁵ parts zinc stearate

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11	1,516,510	11
;	The plastisol thus prepared was lined into the peripheral annular channel of lug caps by means of conventional lining machinery and the lined material was fluxed for	
5	one minute at 350°F (177°C). The resulting gasketed lug caps were tested for sealing ability by determining their vacuum retention performance with the following procedure: A jar is filled with boiling water to one half inch from its top. A lug cap is sealed on with 35 inch pound (0.40 mkg) torque. The resulting assembly is cooled to room temperature and kept	. 5
10	for one week. A vacuum reading is then obtained by means of a vacuum gauge. After removal of the cap, the compressed gasket thickness is measured. All jars tested with lug caps gasketed with the above composition held a vacuum satisfactorily. Gasket thicknesses ranging from 20 to 35 mils (0.5 to 0.9 mm) were acceptable.	10 _.
	Example 34.	
15	A plastisol was prepared with the following ingredients: 100 parts poly(methyl methacrylate), of 35 microns particle size 120 parts acetyl tributyl citrate 5 parts titanium dioxide 5 parts zinc stearate	15
	4 parts azobisformamide	20
20	1.5 parts Lightweight silica (Cab-O-Sil) This composition was lined into a wide-mouth jar closure (83 mm diameter) by conventional means. The lining was fluxed for 1.5 minute at 350°F (177°C). The caps thus obtained were applied to jars with a 32 inch pound (0.36 mkg) torque and kept at room temperature for 24 hours. They were then placed on a	<i>2</i> 0
25	stacker for one week, being subjected in the process to a net load of 128 lbs (37 kg) on each cap and jar assembly. The jars were then removed, kept at room temperature for 24 hours and placed in a cold room for two days. Vacuum was determined at 40°F (4°C) by means of a gauge. Nine out of ten jars tested held 40 mmHg, an accept-	25
	able performance.	20
30	WHAT WE CLAIM IS:— 1. A plastisol comprising (a) 100 weight parts of solid particles of average size 0.1 to 500 microns of an organic polymer consisting substantially of units derived	30
	from (i) one or more of: t-butyl acrylate, a C_{1-4} alkyl methacrylate or cyclohexyl meth-	
35	acrylate or	35
	(ii) one or more of the above monomers and one or more comonomers selected from methacrylates of aliphatic C ₂ to C ₁₀ alcohols, acrylates of aliphatic C ₁ —C ₁₀ alcohols, styrene and alpha-methyl styrene, said polymer having a glass transition temperature of about 35°C and an average degree of polymerisation	40
40	of more than 400, and (b) 30 to 1000 weight parts of an organic liquid component which serves as a compatible plasticiser for the organic polymer, to form a viscous dispersion having a liquid or paste-like appearance and a viscosity index V. (as hereinbefore defined) of less than 3.0.	40
•	2 A plastisol according to claim 1, wherein the organic polymer (a) consists	45
45	substantially of units derived from methyl methacrylate or methyl methacrylate and a comonomer as defined in claim 1. 3. A plastisol according to claim 2, wherein the organic polymer (a) consists substantially of units derived from at least 40% by weight of methyl methacrylate and	75
	substantially all the balance being a C_{2} , alkyl methacrylate or a C_{1-4} alkyl acrylate.	
50	4. A plastisol according to claim 3 wherein the organic polymer (a) contains at least 50% by weight of methyl methacrylate. 5. A plastisol according to claim 1, wherein the organic polymer consists sub-	50
	stantially of units derived from one of:— t-butyl acrylate, ethyl, n-propyl, isopropyl,	
55	n-butyl, sec-butyl, t-butyl, or cyclohexyl methacrylate, or from one of these monomers and a comonomer as defined in claim 1.	55
55	6. A plastisol according to any one of claims 1 to 5 wherein the average particle size of the polymer is related to its glass transition temperature so as to lie within the	
60	triangle XYZ of the accompanying drawing. 7. A plastisol according to claim 6 wherein the plasticiser is dioctyl phthalate, dibutyl phthalate or di(2-methoxyethyl) phthalate and the average particle size of the polymer is related to its glass transition temperature so as to lie within the triangle ABC, DEF or GYH, respectively, of the accompanying drawing.	60

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8. A plastisol according to any one of claims 1 to 7 wherein the organic polymer (a) contains also up to 3% by weight, based on the polymer, of units derived from an acrylic and/or methacrylate monomer having a hydroxyl, epoxy, amino or free carboxyl group.

9. A plastisol according to any one of claims 1 to 7 wherein the organic polymer (a) contains up to 10% by weight of units derived from one or more comonomers containing a heterocyclic group having at least one nitrogen atom in a heterocyclic ring.

10. A plastisol according to claim 9 wherein the proportion of heterocyclic groupcontaining comonomer is from 0.2 to 3.0% by weight.

11. A plastisol according to claim 9 or 10 wherein the heterocyclic group-10 containing comonomer is a heterocyclic vinyl compound of the general formula

$$\begin{array}{c|c}
R_1 \\
 & \downarrow \\
CH_2 = C - (CH_2)_n - R_2
\end{array}$$

in which n is 0 or 1, R₁ is a hydrogen atom, a linear or branched alkyl group of from 1 to 4 carbon atoms or a phenyl group, and R₂ is a group of the formula:

in which R₃ is a heterocyclic group having one or more rings, at least one nitrogen atom being present in at least one ring.

12. A plastisol according to any one of claims 9 to 11, which also contains a crosslinking agent in an amount of from 0.1 to 3.0% by weight which is capable of reacting with the heterocyclic groups of said comonomer.

13. A plastisol according to claim 12, wherein the cross-linking agent is a polyepoxide and/or a polybasic carboxylic acid.

14. A plastisol according to any one of claims 1 to 13 which also contains from

0.1 to 50 parts by weight of an adhesion promoter.

15. A plastisol according to any one of the preceding claims which contains also 25 up to 700 weight parts of inert filler.

16. A plastisol according to claim 2 substantially as described in any one of Examples 3, 4, 6, 8, 10, 15, 16, 17, 18, 19 or 20.

17. A plastisol according to claim 5 substantially as described in any one of

Examples 29 to 32. 18. A plastisol according to claim 1 substantially as described in any one of

Examples 25 to 28, 33 or 34. 19. Shaped plastics articles made by shaping and gelling a plastisol according to

any one of claims 1, 3, 4, 6 to 15 or 18. 20. Shaped plastics articles made by shaping and gelling a plastisol according to

claim 2 or 16. 21. Shaped plastics articles made by shaping and gelling a plastisol according to

claim 5 or 17.

22. A sealing gasket produced by shaping and gelling a plastisol according to any one of claims 1, 3, 4, 6 to 15 or 18.

23. A sealing gasket produced by shaping and gelling a plastisol according to claim 2 or 16.

24. A sealing gasket produced by shaping and gelling a plastisol according to claim 5 or 17.

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1 SHEET

